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TI Separation and Analysis of Plasmid Denatured Forms Using
Hydrophobic Interaction Chromatography

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AB This work explores the possibility of using a hydrophobic
interaction chromatog. (HIC) support to sep. supercoiled plasmids from
denatured forms, by taking advantage of their different surface
hydrophobicity. The hydrophobic gel used in this work
was prep'd. by covalent immobilization of 1,4-butanediol diglycidyl ether
on Sepharose CL-6B (Pharmacia). The hydrophobic interaction
between this support and lipases was previously reported. Expts. were
carried out in a 16 x 150-mm column packed with this gel and equilibrated
with 10 mM Tris, pH 8, with 1.5 M (NH₄)₂SO₄ at a flow rate of 60 mL/h.
The absorbance was monitored at 254 nm. The plasmid used in the expts.
was produced by fermt. of E. coli DH5.alpha. competent cells transformed
with the 8.5-kb pCF1-CFTR plasmid (Genzyme Corp.). Growth was carried

out
overnight in LB medium (30 ug/mL kanamycin), in 100-mL shake-flasks at
37.degree. and 250 rpm. This work shows that HIC can be used for the
sepn. of plasmid variants. The technique can play an important role in
the preparative purifn. of super-coiled plasmids for gene therapy and DNA
vaccination. In fact, the HIC support studied was capable of removing
denatured plasmid variants that are usually produced with the widespread
method of alk. lysis of plasmid isolation. This is very difficult to
achieve using other chromatog. processes. Another important application
could be in the monitoring and quality control of purified plasmids.
Ongoing work indicates also an ability of the HIC support to sep
. RNA and genomic DNA from plasmids. (c)
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